CLAIMS

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What is claimed is:

- 1. An improved method for enhancing immune responses by upregulating co-stimulatory molecules, the upregulating of the co-stimulatory molecules comprising the steps of administering a glucan-containing composition to an animal or a human, in sufficient dosage to cause an enhanced expression of co-stimulatory molecules on antigen presenting cells, the co-stimulatory molecules providing a second signal to T lymphocytes, causing the T lymphocytes to differentiate into armed effector cells.
- The improved method of Claim 1 wherein the glucan-containing composition is at minimum a portion of a glucan selected from the group consisting of β 1,3-glucans and β 1,6glucans.
 - 3. The improved method of Claim 1 wherein the molecule expressed is a molecule from a family of B7 molecules.
- 15 4. The improved method of Claim 5 wherein the family of B7 molecules comprises a molecule selected from the group including B7.1, B7.2, and B7.3.
 - 5. A method for expressing an increased number of B7 molecules on the surface of an antigen presenting cell to more efficiently potentiate the immune system comprising the steps of: obtaining an upregulating agent;
- administering the upregulating agent to an organism; and,
 - allowing an upregulation of B7 molecules on a cell whereby an expression of the B7 molecules allows reaction with an effector cell, the reaction with the armed effector cell potentiating an immune response.

- 6. An enhanced macrophage enhanced by immunological response modification, the macrophage enhancing immunological response, comprising a macrophage enhanced by the delivery of a necessary signal that augments an upregulation of a costimulatory molecule, the enhanced upregulation of the costimulatory molecule, in part, caused by a first glucan containing composition interacting with a second glucan containing composition.
- 7. The macrophage of Claim 6 wherein the costimulatory molecule is a B7 molecule.
- 8. The macrophage of Claim 7 wherein the B7 molecule is selected from a group comprising B7.1, B7.2 and B7.3.
- 9. A beta-glucan preparation which provides a free amino group for conjugation and which can be used as a vaccine adjuvant, comprising:

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microparticulate beta -(1,3)-glucan with or without beta -(1,6)-glucan side chains which do not substantially reaggregate upon drying or rehydration;

about 1-10 % by weight partially deacetylated N-acetylglucosamine within said betaglucan that provides a free amino group for vaccine conjugation; and

- a vaccine or an antigenic substance, wherein said vaccine or antigenic substance is conjugated with said free amino group.
- 10. The preparation of Claim 9, wherein the glucan contains about 1 %-10% by weight chitin or partially deacetylated N-acetylglucosamine.
- 11. A method of using microparticulate beta -(1,3)-glucan as a vaccine adjuvant comprising20 the steps of:

preparing or obtaining a microparticulate beta -(1,3)-glucan composition which does not substantially reaggregate upon drying and rehydration which contains partially deacetylated N-acetlyglucosamine with a free amino group;

suspending the microparticulate beta -(1,3)-glucan composition in liquid; adding at least one vaccine or antigenic substance; conjugating the vaccine onto the free amino group; and administering the vaccine to an animal or human.

- The method of Claim 11, wherein the glucan contains less than 5% by weight protein and lipid, more than 85% by weight glucose, and about 1-10% by weight chitin or partially deacetylated N-acetylglucosamine.
- 13. A vaccine adjuvant which contains microparticulate beta glucan with a
 free amino group, which enhances the immunologic effects of vaccine or antigenic substance,
 comprising:

microparticulate beta -(1,3)-glucan with or without beta -(1,6)-glucan side chains which do not substantially reaggregate upon drying or rehydration;

at least 2% by weight partially deacetylated N-acetylglucosamine within said beta-glucan that provides a free amino group for vaccine conjugation.

14. A vaccine conjugate or conjugated antigenic substance attached to the free amino group of microparticulate beta -(1,3)-glucan, which stabilizes the vaccine and enhances the immunologic effects of vaccine, comprising:

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microparticulate beta -(1,3)-glucan with or without beta -(1,6)-glucan side chains with about 1-10% by weight partially deacetylated N-acetylglucosamine within said beta-glucan that provides a free amino group for vaccine conjugation which does not substantially reaggregate upon drying or rehydration;

a vaccine or an antigenic substance, wherein said vaccine or antigenic substance is conjugated with said free amino group.

15. A method for preparing a small particle size glucan for dry packaging comprising the steps of:

obtaining a polysaccharide composition comprising the glucan;

hydrating the glucan with a liquid;

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loading the glucan in a sprayer; and,

spraying the glucan.

16. The method of Claim 15 further comprising the steps of:

grinding the glucan and

- 10 re-hydrating the glucan whereby a portion of the glucan is dissociated into particles of about 1 2 microns in diameter.
 - 17. The method of Claim 15 wherein the glucan is substantially glucan selected from the group comprising beta-(1,3)-glucan and beta-(1,6)-glucan.
 - 18. The method of Claim 15 wherein the disrupting is accomplished by sonicating the glucan.
 - 19. A method for preparing a small particle size glucan for improved immunological response through enhanced activation of a macrophages and freeze drying the glucan such that re-hydration of the glucan disassociates the glucan, comprising the steps of:

obtaining a polysaccharide composition comprising a glucan containing composition;

20 hydrating the glucan containing composition with a liquid;

disrupting the glucan;

adding a gelatin solution to the hydrated glucan; and,

freeze drying the glucan.

- 20. The method of Claim 19 further comprising the step of grinding the glucan.
- 21. The method of Claim 19 further comprising the step of rehydrating the glucan whereby a portion of the glucan is dissociated into particles of .3 3.0 microns in diameter.
- 22. The method of Claim 19, wherein the disrupting is accomplished by sonicating the glucan.
- 23. The method of Claim 19, wherein the glucan is substantially glucan selected from the group comprising beta-(1,3)-glucan and beta-(1,6)-glucan.

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